CLAIMS

What is claimed is:

- 1. A method to select altered peptide species for administration to a subject, said subject presenting a native ligand for which activation of an immune response against said native ligand is desired, comprising the steps of:
- a. identifying a plurality of altered peptide species capable of eliciting
 an immune response to said native ligand;
- b. selecting at least two altered peptide species wherein each altered peptide activates a population of T cells having a distinct T cell receptor $V\beta$ recombination.
- 2. The method of claim 1, further comprising the step of administering said selected altered peptide species to said subject.
- 3. A method to select altered peptide species for administration to a member of a population of subjects having a given HLA-type, said population presenting a native ligand for which activation of an immune response against said native ligand is desired, comprising the steps of:
- a. identifying a plurality of altered peptide species capable of eliciting an immune response to said native ligand; and
- b. selecting two or more of said altered peptide species, wherein each of said altered peptide species activates a T cell having a distinct T cell receptor $V\beta$ recombination in a sample representative of said population.
- 4. The method of claim 3, further comprising the step of administering said selected altered peptide species to said member.
 - 5. The method of claim 3, wherein said HLA-type is HLA-A2.

- 6. The method of claims 1 or 3, comprising from at least 2 to 6 different altered peptide species.
- 7. The method of claims 1 or 3, comprising at least 3 different altered peptide species.
- 8. The method of claims 1 or 3, wherein at least one altered peptide species is covalently linked to one or more amino acids naturally contiguous to said native human ligand.
- 9. The method of claims 1 or 3, wherein said native ligand is a mammalian tumor epitope.
- 10. The method of claims 1 or 3, wherein said native ligand is a human viral antigen.
- 11. The method of claims 1 or 3, further comprising the step of formulating said selected altered peptides in a pharmaceutically acceptable carrier suitable for administration to humans.
- 12. A composition comprising multiple peptide ligand species directed at a single native ligand, wherein at least two altered peptide species activate a different T cell clone from each other and the T cell receptor $V\beta$ recombination of each of said activated T cell clones is different.
- 13. A composition comprising two or more altered peptide species, wherein each of said two or more species is characterized by an ability to activate a different subpopulation of cytotoxic T lymphocytes (CTLs) against the same native ligand.

- 14. The composition of claims 12 or 13, comprising from at least 2 to 6 different peptide species.
- 15. The composition of claims 12 or 13, comprising at least 3 different peptide species.
- 16. The composition of claims 12 or 13, wherein at least one peptide species is covalently linked to one or more amino acids naturally contiguous to said native human ligand.
- 17. The composition of claim 13, wherein said altered peptide species activate said different subpopulations in one member of a selected population of subjects
- 18. The composition of claim 13, wherein said altered peptide species activate a different subpopulation in two or more members of said population of subjects.
- 19. The composition of claims 12 or 13, wherein said native ligand is a mammalian tumor epitope.
- 20. The composition of claims 12 or 13, wherein said native ligand is a human viral antigen.
- 21. The composition of claims 12 or 13, further comprising a pharmaceutically acceptable carrier.
 - 22. A kit comprising:
 - a. multiple peptide species directed at a single native ligand, wherein
 - i. a first peptide species activates a first T cell,

- ii. a second peptide species activates a second T cell, and the T cell receptor V β recombination of said first T cell is different from the T cell receptor V β recombination of said second T cell; and
 - b. instructions for the co-administration of each of said peptides.
- 23. The kit of claim 22, wherein said selected peptide species are packaged in combination.
- 24. The kit of claim 22, further comprising instructions to identify subjects who exhibit a positive therapeutic response to the administration of said multiple peptide ligand species.
 - 25. A method comprising the steps of:
- a. identifying a plurality of altered peptide species characterized by an ability to elicit an immune response against the same native ligand;
- b. determining the T cell receptor $V\beta$ recombination profile of the T cell population activated by each of said identified altered peptide species in a plurality of test subjects; and

selecting at least two or more altered peptide species wherein each of said altered peptide species activates T cell populations having distinct T cell receptor $V\beta$ recombinations in a majority of said test subjects.

- 26. The method of claim 25, wherein at least one of said selected altered peptide species is more than said native antigen.
- 27. The method of claim 25, further comprising the step of administering said selected altered peptide species to a subject having said native antigen.

- 28. The method of claim 25, further comprising the step of packaging said selected altered peptide species in a form suitable for administration to a subject.
- 29. The method of claim 26, wherein said selected altered peptide species are packaged together.